

Roberto VILLA et al.

IN THE ABSTRACT:

Please delete the abstract as originally filed which appears on the cover sheet of the Published Application.  
~~Add~~ new abstract as enclosed herewith on a separate sheet.

REMARKS

Claims 3, 6 and 8-11 were amended to correct multiple dependency. Attached hereto is a marked-up version of the changes made to the claims by the current amendment. The attached page is captioned "VERSION WITH MARKINGS TO SHOW CHANGES MADE".

SCANNED, # 12

Respectfully submitted,

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**"VERSION WITH MARKINGS TO SHOW CHANGES MADE"**

Claims 3, 6 and 8-11 have been amended as follows:

3. **(Amended)** Compositions as claimed in claim 1-~~or~~-2, wherein 5-aminosalicylic acid is inglobated in the molten lipophilic matrix by kneading, extrusion and/or granulation.

6. **(Amended)** Compositions as claimed in ~~any one of the above claims, claim 1,~~ comprising a gastro-resistant outer coating.

8. **(Amended)** Compositions as claimed in ~~any one of the above claims, claim 1,~~ in the form of tablets, capsules, minitablets, wherein the active ingredient is completely contained inside the lipophilic matrix.

9. **(Amended)** Compositions as claimed in ~~any one of claims 1 to 7, claim 1,~~ in the form of tablets, capsules, minitablets, wherein the active ingredient is dispersed both in the hydrophilic matrix and the lipophilic matrix.

10. **(Amended)** Compositions as claimed in ~~any one of the above claims, claim 1,~~ wherein the percentage of the active ingredient on the total composition weight ranges from 80 to 95%.

11. **(Amended)** A process for the preparation of the compositions of ~~claims 1-10, claim 1,~~ which comprises:

- a) melt granulation of at least one portion of the active ingredient with the lipophilic excipients with melting point lower than 90°C;
- b) mixing the granules from step a) with the hydrophilic excipients and subsequent tabletting or compression.